Under the Pasarwork Reduction Act of 1995, to earsons are required to respond to a collection of information unless it disclays a valid UME control number.

Under the Passerver's Recyclic Access REQUEST FOR ACCESS	1995, 100 OSTICOTA BEST ESSUÉRAS POPES TO AN ABANDONE	D APPLICATION U	NDER 37 CFR 1.14
KERNER LOK WOOFGO	10 A.C. 121	in re Application of	<i>(</i>
-		Queer e	
Bring completed form to: File Information Unit Oristal Pieze Three, Room 1001		07/310, 2 50	2/13/89
07/31 2021 South Clarx Place Arlington, VA Telephone: (703) 306-2733			Papar No. 54 }
			(
l hereby request access under 37 application, which is identified in attachment):	OFR $1.14(z)(1)(iv)$ to the sq , or to which z benefit is dis	ocileation file record of the imed, in the following de	he shove-identified ABANDONED soument (as shown in the
Matter Steine Gatest 450	lication Publication No	, page,	line
Grades Clates Catest Mus	niber <u>5,530,101</u> , co	lums, line,	Gf Gf
			•
WIFO Pub. No	, page		
For unpublished applications (1) If the benefit of the pend application that has: (a) patent application public Article 21(2), a member the file contents; the pending application is inco- registration, a U.S. pate	or the pending epindenent that are sail pending:	ent application publication pay of: ation. arrivisa identified in a U.S. p	in accordance with PC1 patent, a statutory invention plication publication in
Rignature		<u> 7/11/65</u>	RECEIVED
Typed or printed na	ime		JUL T 1005
Registration Number	if applicable	Andrew E	ile Information Unit]
(703) 521-1952		Unit: _	
Telechone Nu		!	

This collection of information is required by 37 OFR 1.14. The information is required to obtain or retain a benefit by the public which is to file (and by the Eustra of the process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 OFR 1.14. This collection is estimated to take 13 minutes to complete, including the process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 OFR 1.14. This collection is estimated to take 13 minutes to complete. Any comments on the estimation, preparing, and submitting the completed application form to the USFTO. Time will vary depending upon the individual case. Any comments on the estimation, preparing, and submitting the complete despite application successful of the USFTO. Time will vary depending upon the individual case. Any comments on the estimation of the Ohier information Office, U.S. Paperment of Commerce, P.O. Sox 1450, Alexandria, VA. 2013;1450, DO NOT SENO FEES OR COMPLETED FORMS TO THIS Information Unit. Crystal Plaza Three. Room 1001, 2021 South Clark Place. Adjunction VA. ADDRESS, BRING TO: File Information Unit. Crystal Plaza Three. Room 1001, 2021 South Clark Place. Adjunction Unit. ADDRESS. BRING TO: File Information Unit, Crystal Plaza Three, Room 1001, 2021 South Clark Place, Arlington, VA.



United States Patent [19]

Queen et al.

[11] Patent Number:

5,530,101

[45] Date of Patent:

Jun. 25, 1996

[54] HUMANIZED IMMUNOGLOBULINS

- [75] Inventors: Cary L. Queen, Los Altos; Harold E. Selick, Belmont, both of Calif.
- [73] Assignee: Protein Design Labs, Inc., Mountain View, Calif.
- [21] Appl. No.: 634,278
- [22] Filed: Dec. 19, 1990

Related U.S. Application Data

- [63] Continuation-in-part of Ser. No. 590,274, Sep. 28, 1990, abandoned, and a continuation-in-part of Ser. No. 310,252; Feb. 13, 1989, abandoned, which is a continuation-in-part of Ser. No. 290,975, Dec. 28, 1988, abandoned.
- [51] Int. Cl.⁶ A61K 39/395; C07K 16/28

[56] References Cited

U.S. PATENT DOCUMENTS

4,816,397	3/1989	Boss et al 435/68
4,816,567	3/1989	Cabilly et al 530/387
4,867,973	9/1989	Geers et al
5,225,539	7/1993	Winter.

FOREIGN PATENT DOCUMENTS

0171496	2/1986	European Pat. Off
0173494	3/1986	European Pat. Off
0184187	6/1986	European Pat. Off
0239400	9/1987	European Pat. Off
0266663	6/1988	European Pat. Off
2188941	10/1987	United Kingdom .
WO86/05513	9/1986	WIPO .
WO87/02671	5/1987	WIPO .
WO89/01783	3/1989	WIPO .

OTHER PUBLICATIONS

Vitteta et al., "Redesigning Nature's Poisons to Create Anti-Tumor Reagents," Science 238:1098-1104 (1987). Ellison et al., "The nucleotide sequence of a human immunoglobulin C(gamma)₁ gene", Nucleic Acids Res. 10:4071-(1982).

Hieter et al., "Cloned Human and Mouse Kappa Immunoglobulin Constatn and J Region Genes Conserve homology in Functional Segments", Cell 22:197-207 (1980).

Sharon et al., "Expression of a V_rC_r chimaeric protein in mouse myeloma cells", *Nature* 309:364-367 (1984).

Takeda et al., "Construction of chimaeric processed immunoglobulin genes containing mouse variable and human constant region sequences", Nature 314:452-454 (1985).

Tan et al., "A Human-Mouse Chimeric Immunoglobulin Gene with a Human Variable Region is Expressed in Mouse Myeloma Cells", *J. Immunol.* 135:3564-3567 (1985).

Morrison et al., "Chimeric human antibody molecules: Mouse antigen-binding domains with human constant region domains," *Proc. Natl. Acad. Sci.* 81:6851-6859 (1984).

Boulianne et al., "Production of functional chimeric mouse/ human antibody," *Nature* 312:643-646 (1984). Neuberger et al., "A hapten-specific chimeric IgE antibody with human physiological effector function," *Nature* 314:268-270 (1985).

Morrison, S. L., "Transfectomas Provide Novel Chimeric Antibodies," *Science* 229:1202-1207 (1985).

Sahagan et al., "A Genetically Engineered Murine/Human Chimeric Antibody Retains Specificity for Human Tumor-Associated Antigen", J. Immunol. 137:1066-1074 (1986). Liu et al., "Expression of mouse::human immunoglobulin heavy-chain cDNA in lymphoid cells", Gene 54:33-40 (1987).

Better et al., "Escherichia coli Secretion of an Active Chimeric Antibody Fragment", Science 240:1041-1043 (1988).

Waldmann, T. A., "The Structure, Function, and Expression of Interleukin-2 Receptors on Normal and Malignant Lymphocytes," *Science* 232:727-732 (1986).

Leonard et al., "The human receptor for T-cell growth factor," J. Biol. Chem. 260:1872-1880 (1985).

Farrar, J., "The biochemistry, biology, and role of interleukin-2 in the induction of cytotoxic T cell and antibodyforming B cell receptors," *Immunol. Rev.* 63:129-166 (1982).

Greene et al., "Growth of Human T Lymphocytes: An Analysis of Interleukin 2 and Its Cellular receptor", in *Progress in Hematology XIV*, E. Brown ed., Grune and Statton, New York (1986) pp. 283-301.

Verhoyen et al., "Reshaping Human Antibodies: Grafting an Antilysozyme Activity", Science 239:1534-1536 (1988).

Jones et al., "Replacing the complementarity-determining regions in a human antibody with those from a mouse", *Nature* 321:522-525 (1986).

Hale et al., "Remission Induction in Non-Hodgkin Lymphoma with Reshaped Human Monoclonal Antibody CAMPATH-1H", Lancet Dec. 17, 1988, pp. 1394-1399. Chothia, C. and A. M. Lesk, "Canonical Structures for the Hypervariable Regions of Immunoglobulins", J. Mol. Biol. 196:901-917 (1987).

(List continued on next page.)

Primary Examiner—Lila Feisee
Attorney, Agent, or Firm—Townsend and Townsend and
Crew

[57] ABSTRACT

Novel methods for producing, and compositions of, humanized immunoglobulins having one or more complementarity determining regions (CDR's) and possible additional amino acids from a donor immunoglobulin and a framework region from an accepting human immunoglobulin are provided. Each humanized immunoglobulin chain will usually comprise, in addition to the CDR's, amino acids from the donor immunoglobulin framework that are, e.g., capable of interacting with the CDR's to effect binding affinity, such as one or more amino acids which are immediately adjacent to a CDR in the donor immunoglobulin or those within about 3 A as predicted by molecular modeling. The heavy and light chains may each be designed by using any one or all of various position criteria. When combined into an intact antibody, the humanized immunoglobulins of the present invention will be substantially non-immunogenic in humans and retain substantially the same affinity as the donor immunoglobulin to the antigen, such as a protein or other compound containing an epitope.

13 Claims, 55 Drawing Sheets

54